

OIPE

RAW SEQUENCE LISTING

SEQUENCE LISTING

DATE: 56/14/2001

PATENT APPLICATION: US/09/841,553

TIME: 12:35:12

Input Set : A:\09841553.txt

Output Set: N:\CRF3\06142001\I841553.raw

ENTERED

```
3 (1) GENERAL INFORMATION:
             (i) APPLICANT: TAKAKURA, Hikaru
     5
                            MORISHITA, Mio
     6
                            YAMAMOTO, Katsuhiko
     7
                            MITTA, Masanori
     8
                            ASADA, Kiyozo
     9
                            TSUNASAWA, Susumu
    10
                             KATO, Ikunoshin
    11
            (ii) TITLE OF INVENTION: HYPERTHERMOSTABLE PROTEASE GENES
    13
           (iii) NUMBER OF SEQUENCES: √42
    15
            (iv) CORRESPONDENCE ADDRESS:
    17
                  (A) ADDRESSEE: Browdy and Neimark
    18
                  (B) STREET: 419 Seventh Street N.W., Ste. 300
    19
    20
                  (C) CITY: Washington
                   (D) STATE: D.C.
    21
                  (E) COUNTRY: United States of America
    22
                  (F) ZIP: 20004
     23
             (v) COMPUTER READABLE FORM:
     25
                   (A) MEDIUM TYPE: Floppy disk
     26
                   (B) COMPUTER: IBM PC compatible
     27
                   (C) OPERATING SYSTEM: PC-DOS/MS-DOS
     28
                   (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
     29
            (vi) CURRENT APPLICATION DATA:
     31
                   (A) APPLICATION NUMBER: US/09/841,553
C--> 32
                   (B) FILING DATE: 24-Apr-2001
C--> 33
           (vii) PRIOR APPLICATION DATA:
C--> 39
                   (A) APPLICATION NUMBER: US 08/894,818
     36
                   (B) FILING DATE:
     37
                   (A) APPLICATION NUMBER: JP 323285/1995
     40
                   (B) FILING DATE: 12-DEC-1995
     41
          (viii) ATTORNEY/AGENT INFORMATION:
     43
                   (A) NAME: Browdy, Roger L.
     44
                   (B) REGISTRATION NUMBER: 25,618
     45
                   (C) REFERENCE/DOCKET NUMBER: TAKAKURA=1
     46
            (ix) TELECOMMUNICATION INFORMATION:
     48
                   (A) TELEPHONE: (202) 628-5197
     49
                   (B) TELEFAX: (202) 737-3528
     50
     53 (2) INFORMATION FOR SEQ ID NO: 1:
             (i) SEQUENCE CHARACTERISTICS:
     55
                   (A) LENGTH: 659 amino acids
     56
                   (B) TYPE: amino acid
     57
                   (C) STRANDEDNESS: single
     58
                   (D) TOPOLOGY: linear
     59
             (ii) MOLECULE TYPE: peptide
     61
             (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
     65 Met Lys Arg Leu Gly Ala Val Val Leu Ala Leu Val Leu Val Gly
```

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Input Set : A:\09841553.txt
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```
68 Leu Leu Ala Gly Thr Ala Leu Ala Ala Pro Val Lys Pro Val Val
66
                                         25
71 Arg Asn Asn Ala Val Gln Gln Lys Asn Tyr Gly Leu Leu Thr Pro
69
                                         40
74 Gly Leu Phe Lys Lys Val Gln Arg Met Asn Trp Asn Gln Glu Val
                                         55
                    50
77 Asp Thr Val Ile Met Phe Gly Ser Tyr Gly Asp Arg Asp Arg Ala
                     65
80 Val Lys Val Leu Arg Leu Met Gly Ala Gln Val Lys Tyr Ser Tyr
                                         85
                     80
83 Lys Ile Ile Pro Ala Val Ala Val Lys Ile Lys Ala Arg Asp Leu
                                        100
                     95
86 Leu Leu Ile Ala Gly Met Ile Asp Thr Gly Tyr Phe Gly Asn Thr
                                        115
                    110
89 Arg Val Ser Gly Ile Lys Phe Ile Gln Glu Asp Tyr Lys Val Gln
                                        130
                    125
92 Val Asp Asp Ala Thr Ser Val Ser Gln Ile Gly Ala Asp Thr Val
                                        145
                    140
95 Trp Asn Ser Leu Gly Tyr Asp Gly Ser Gly Val Val Val Ala Ile
                                        160
                    155
98 Val Asp Thr Gly Ile Asp Ala Asn His Pro Asp Leu Lys Gly Lys
                                         175
                    170
 101 Val Ile Gly Trp Tyr Asp Ala Val Asn Gly Arg Ser Thr Pro Tyr
                                          190
                     185
 104 Asp Asp Gln Gly His Gly Thr His Val Ala Gly Ile Val Ala Gly
                     200
 107 Thr Gly Ser Val Asn Ser Gln Tyr Ile Gly Val Ala Pro Gly Ala
                                          220
                     215
 110 Lys Leu Val Gly Val Lys Val Leu Gly Ala Asp Gly Ser Gly Ser
                                          235
                     230
 113 Val Ser Thr Ile Ile Ala Gly Val Asp Trp Val Val Gln Asn Lys
                                          250
                      245
 116 Asp Lys Tyr Gly Ile Arg Val Ile Asn Leu Ser Leu Gly Ser Ser
                                                               270
                                          265
                      260
 119 Gln Ser Ser Asp Gly Thr Asp Ser Leu Ser Gln Ala Val Asn Asn
                                          280
                      275
 122 Ala Trp Asp Ala Gly Ile Val Val Cys Val Ala Ala Gly Asn Ser
                                           295
                      290
 125 Gly Pro Asn Thr Tyr Thr Val Gly Ser Pro Ala Ala Ala Ser Lys
                                           310
                      305
 128 Val Ile Thr Val Gly Ala Val Asp Ser Asn Asp Asn Ile Ala Ser
                                           325
                      320
 131 Phe Ser Ser Arg Gly Pro Thr Ala Asp Gly Arg Leu Lys Pro Glu
                                           340
                      335
 135 Val Val Ala Pro Gly Val Asp Ile Ile Ala Pro Arg Ala Ser Gly
                      350
  138 Thr Ser Met Gly Thr Pro Ile Asn Asp Tyr Tyr Thr Lys Ala Ser
                                           370
                       365
  139
```

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```
141 Gly Thr Ser Met Ala Thr Pro His Val Ser Gly Val Gly Ala Leu
                                         385
                    380
144 Ile Leu Gln Ala His Pro Ser Trp Thr Pro Asp Lys Val Lys Thr
                                         400
                    395
147 Ala Leu Ile Glu Thr Ala Asp Ile Val Ala Pro Lys Glu Ile Ala
                                         415
148
150 Asp Ile Ala Tyr Gly Ala Gly Arg Val Asn Val Tyr Lys Ala Ile
                    425
151
153 Lys Tyr Asp Asp Tyr Ala Lys Leu Thr Phe Thr Gly Ser Val Ala
                                         445
                    440
156 Asp Lys Gly Ser Ala Thr His Thr Phe Asp Val Ser Gly Ala Thr
                                         460
                    455
159 Phe Val Thr Ala Thr Leu Tyr Trp Asp Thr Gly Ser Ser Asp Ile
                                         475
                    470
162 Asp Leu Tyr Leu Tyr Asp Pro Asn Gly Asn Glu Val Asp Tyr Ser
                                         490
                    485
165 Tyr Thr Ala Tyr Tyr Gly Phe Glu Lys Val Gly Tyr Tyr Asn Pro
                                         505
                    500
168 Thr Ala Gly Thr Trp Thr Val Lys Val Val Ser Tyr Lys Gly Ala
                                         520
                    515
171 Ala Asn Tyr Gln Val Asp Val Val Ser Asp Gly Ser Leu Ser Gln
                                         535
                    530
172
174 Ser Gly Gly Asn Pro Asn Pro Asn Pro Asn Pro Asn Pro Thr
                                         550
                     545
175
177 Pro Thr Thr Asp Thr Gln Thr Phe Thr Gly Ser Val Asn Asp Tyr
                                         565
                     560
178
180 Trp Asp Thr Ser Asp Thr Phe. Thr Met Asn Val Asn Ser Gly Ala
                                         580
                     575
183 Thr Lys Ile Thr Gly Asp Leu Thr Phe Asp Thr Ser Tyr Asn Asp
                                         595
                     590
186 Leu Asp Leu Tyr Leu Tyr Asp Pro Asn Gly Asn Leu Val Asp Arg
                                                              615
                                         610
                     605
187
189 Ser Thr Ser Ser Asn Ser Tyr Glu His Val Glu Tyr Ala Asn Pro
                                         625
                     620
192 Ala Pro Gly Thr Trp Thr Phe Leu Val Tyr Ala Tyr Ser Thr Tyr
                                         640
                                                              645
                     635
195 Gly Trp Ala Asp Tyr Gln Leu Lys Ala Val Val Tyr Tyr Gly
                                         655
                     650
196
199 (2) INFORMATION FOR SEQ ID NO: 2:
          (i) SEQUENCE CHARACTERISTICS:
201
               (A) LENGTH: 1977 base pairs
202
               (B) TYPE: nucleic acid
203
               (C) STRANDEDNESS: double
204
               (D) TOPOLOGY: linear
205
         (ii) MOLECULE TYPE: genomic DNA
207
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
211 ATGAAGAGGT TAGGTGCTGT GGTGCTGGCA CTGGTGCTCG TGGGTCTTCT GGCCGGAACG
                                                                          60
213 GCCCTTGCGG CACCCGTAAA ACCGGTTGTC AGGAACAACG CGGTTCAGCA GAAGAACTAC
                                                                          120
215 GGACTGCTGA CCCCGGGACT GTTCAAGAAA GTCCAGAGGA TGAACTGGAA CCAGGAAGTG
                                                                          180
```

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```
217 GACACCGTCA TAATGTTCGG GAGCTACGGA GACAGGGACA GGGCGGTTAA GGTACTGAGG
                                                                          240
    219 CTCATGGGCG CCCAGGTCAA GTACTCCTAC AAGATAATCC CTGCTGTCGC GGTTAAAATA
                                                                           300
    221 AAGGCCAGGG ACCTTCTGCT GATCGCGGGC ATGATAGACA CGGGTTACTT CGGTAACACA
                                                                           360
    223 AGGGTCTCGG GCATAAAGTT CATACAGGAG GATTACAAGG TTCAGGTTGA CGACGCCACT
    225 TCCGTCTCCC AGATAGGGGC CGATACCGTC TGGAACTCCC TCGGCTACGA CGGAAGCGGT
                                                                           480
    227 GTGGTGGTTG CCATCGTCGA TACGGGTATA GACGCGAACC ACCCCGATCT GAAGGGCAAG
                                                                           540
    229 GTCATAGGCT GGTACGACGC CGTCAACGGC AGGTCGACCC CCTACGATGA CCAGGGACAC
    231 GGAACCCACG TTGCGGGTAT CGTTGCCGGA ACCGGCAGCG TTAACTCCCA GTACATAGGC
    233 GTCGCCCCCG GCGCGAAGCT CGTCGGCGTC AAGGTTCTCG GTGCCGACGG TTCGGGAAGC
                                                                           720
    235 GTCTCCACCA TCATCGCGGG TGTTGACTGG GTCGTCCAGA ACAAGGACAA GTACGGGATA
    237 AGGGTCATCA ACCTCTCCCT CGGCTCCTCC CAGAGCTCCG ACGGAACCGA CTCCCTCAGT
    239 CAGGCCGTCA ACAACGCCTG GGACGCCGGT ATAGTAGTCT GCGTCGCCGC CGGCAACAGC
    241 GGGCCGAACA CCTACACCGT CGGCTCACCC GCCGCCGCGA GCAAGGTCAT AACCGTCGGT
    243 GCAGTTGACA GCAACGACAA CATCGCCAGC TTCTCCAGCA GGGGACCGAC CGCGGACGGA
                                                                          1020
    245 AGGCTCAAGC CGGAAGTCGT CGCCCCCGGC GTTGACATCA TAGCCCCGCG CGCCAGCGGA
                                                                          1080
    247 ACCAGCATGG GCACCCCGAT AAACGACTAC TACACCAAGG CCTCTGGAAC CAGCATGGCC
                                                                          1140
    249 ACCCCGCACG TTTCGGGCGT TGGCGCGCTC ATCCTCCAGG CCCACCCGAG CTGGACCCCG
                                                                          1200
    251 GACAAGGTGA AGACCGCCCT CATCGAGACC GCCGACATAG TCGCCCCCAA GGAGATAGCG
                                                                          1260
    253 GACATCGCCT ACGGTGCGGG TAGGGTGAAC GTCTACAAGG CCATCAAGTA CGACGACTAC
    255 GCCAAGCTCA CCTTCACCGG CTCCGTCGCC GACAAGGGAA GCGCCACCCA CACCTTCGAC
    257 GTCAGCGGCG CCACCTTCGT GACCGCCACC CTCTACTGGG ACACGGGCTC GAGCGACATC
                                                                          1440
    259 GACCTCTACC TCTACGACCC CAACGGGAAC GAGGTTGACT ACTCCTACAC CGCCTACTAC
                                                                          1500
    261 GGCTTCGAGA AGGTCGGCTA CTACAACCCG ACCGCCGGAA CCTGGACGGT CAAGGTCGTC
                                                                          1560
    263 AGCTACAAGG GCGCGGCGAA CTACCAGGTC GACGTCGTCA GCGACGGGAG CCTCAGCCAG
    265 TCCGGCGGCG GCAACCCGAA TCCAAACCCC AACCCGAACC CAACCCCGAC CACCGACACC
    267 CAGACCTTCA CCGGTTCCGT TAACGACTAC TGGGACACCA GCGACACCTT CACCATGAAC
    271 CTCGACCTCT ACCTCTACGA CCCCAACGGC AACCTCGTTG ACAGGTCCAC GTCGAGCAAC
    273 AGCTACGAGC ACGTCGAGTA CGCCAACCCC GCCCCGGGAA CCTGGACGTT CCTCGTCTAC
                                                                         1920
    275 GCCTACAGCA CCTACGGCTG GGCGGACTAC CAGCTCAAGG CCGTCGTCTA CTACGGG
                                                                          1977
    278 (2) INFORMATION FOR SEQ ID NO: 3:
             (i) SEQUENCE CHARACTERISTICS:
    280
                  (A) LENGTH: 522 amino acids
    281
                  (B) TYPE: amino acid
    282
                  (C) STRANDEDNESS: single
    283
                  (D) TOPOLOGY: linear
    284
            (ii) MOLECULE TYPE: peptide
     286
            (ix) FEATURE:
C--> 288
                 OTHER INFORMATION:
W--> 289
W--> 290 Xaa at position 428 is Gly or Val.
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
     293 Ala Glu Leu Glu Gly Leu Asp Glu Ser Ala Ala Gln Val Met Ala
                                             10
     296 Thr Tyr Val Trp Asn Leu Gly Tyr Asp Gly Ser Gly Ile Thr Ile
                                            25
                         20
     299 Gly Ile Ile Asp Thr Gly Ile Asp Ala Ser His Pro Asp Leu Gln
                                40
                         35
     302 Gly Lys Val Ile Gly Trp Val Asp Phe Val Asn Gly Arg Ser Tyr
                         50
     303
```

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```
305 Pro Tyr Asp Asp His Gly His Gly Thr His Val Ala Ser Ile Ala
                                            70
                       65
  308 Ala Gly Thr Gly Ala Ala Ser Asn Gly Lys Tyr Lys Gly Met Ala
                                            85
                       80
  309
  311 Pro Gly Ala Lys Leu Ala Gly Ile Lys Val Leu Gly Ala Asp Gly
                                           100
                        95
  312
  314 Ser Gly Ser Ile Ser Thr Ile Ile Lys Gly Val Glu Trp Ala Val
                                           115
  317 Asp Asn Lys Asp Lys Tyr Gly Ile Lys Val Ile Asn Leu Ser Leu
                                           130
                       125
  320 Gly Ser Ser Gln Ser Ser Asp Gly Thr Asp Ala Leu Ser Gln Ala
                                           145
                       140
  323 Val Asn Ala Ala Trp Asp Ala Gly Leu Val Val Val Ala Ala
                                           160
                       155
  326 Gly Asn Ser Gly Pro Asn Lys Tyr Thr Ile Gly Ser Pro Ala Ala
                                           175
                       170
  329 Ala Ser Lys Val Ile Thr Val Gly Ala Val Asp Lys Tyr Asp Val
                                           190
                       185
  332 Ile Thr Ser Phe Ser Ser Arg Gly Pro Thr Ala Asp Gly Arg Leu
                                           205
                       200
  336 Lys Pro Glu Val Val Ala Pro Gly Asn Trp Ile Ile Ala Ala Arg
                                                                225
                                           220
                       215
   337
  339 Ala Ser Gly Thr Ser Met Gly Gln Pro Ile Asn Asp Tyr Tyr Thr
   340
   342 Ala Ala Pro Gly Thr Ser Met Ala Thr Pro His Val Ala Gly Ile
                                            250
                       245
   345 Ala Ala Leu Leu Gln Ala His Pro Ser Trp Thr Pro Asp Lys
                       260
   346
   348 Val Lys Thr Ala Leu Ile Glu Thr Ala Asp Ile Val Lys Pro Asp
                                            280
                       275
   349
   351 Glu Ile Ala Asp Ile Ala Tyr Gly Ala Gly Arg Val Asn Ala Tyr
                                            295
                       290
   354 Lys Ala Ile Asn Tyr Asp Asn Tyr Ala Lys Leu Val Phe Thr Gly
                                                                315
                                            310
                       305
   357 Tyr Val Ala Asn Lys Gly Ser Gln Thr His Gln Phe Val Ile Ser
                                            325
                       320
   360 Gly Ala Ser Phe Val Thr Ala Thr Leu Tyr Trp Asp Asn Ala Asn
                                            340
                       335
   363 Ser Asp Leu Asp Leu Tyr Leu Tyr Asp Pro Asn Gly Asn Gln Val
                                            355
                        350
   366 Asp Tyr Ser Tyr Thr Ala Tyr Tyr Gly Phe Glu Lys Val Gly Tyr
                                                                 375
                                            370
                        365
   369 Tyr Asn Pro Thr Asp Gly Thr Trp Thr Ile Lys Val Val Ser Tyr
                                            385
                        380
   372 Ser Gly Ser Ala Asn Tyr Gln Val Asp Val Val Ser Asp Gly Ser
                                            400
                        395
   375 Leu Ser Gln Pro Gly Ser Ser Pro Ser Pro Gln Pro Glu Pro Thr
                        410
                                            415
--> 378 Val Asp Ala Lys Thr Phe Gln Xaa Ser Asp His Tyr Tyr Tyr Asp
```

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/841,553

DATE: 06/14/2001 TIME: 12:35:13

Input Set : A:\09841553.txt

Output Set: N:\CRF3\06142001\I841553.raw

L:414 M:246 W: Invalid value of Alpha Sequence Header Field, [FEATURE:], SeqNo=4

L:32 M:220 C: Keyword misspelled or invalid format, [(A) APPLICATION NUMBER:]
L:33 M:220 C: Keyword misspelled or invalid format, [(B) FILING DATE:]
L:35 M:220 C: Keyword misspelled or invalid format, [(vii) PRIOR APPLICATION DATA:]
L:39 M:220 C: Keyword misspelled or invalid format, [(vii) PRIOR APPLICATION DATA:]
L:288 M:220 C: Keyword misspelled or invalid format, [(ix) FEATURE:]
L:289 M:246 W: Invalid value of Alpha Sequence Header Field, Data=[Xaa at position 428 is Gly or Val.]
L:291 M:246 W: Invalid value of Alpha Sequence Header Field, [FEATURE:], SeqNo=3
L:378 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:3
L:411 M:220 C: Keyword misspelled or invalid format, [(ix) FEATURE:]
L:412 M:246 W: Invalid value of Alpha Sequence Header Field, Data=[N at position 1283 is G or T.]